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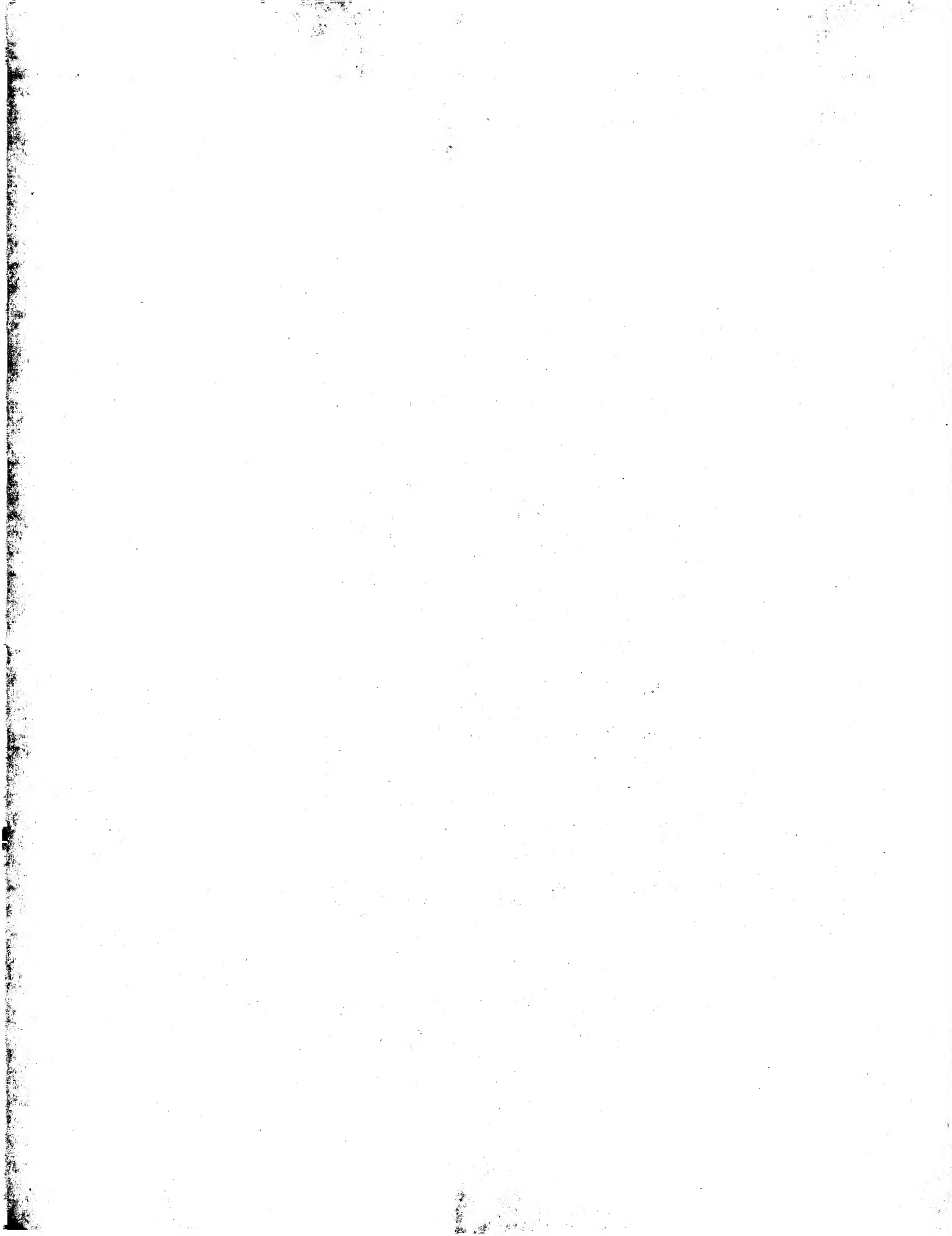
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Hakija  
Applicant

1. Lilius, Esa-Matti  
Kaarina
2. Virta, Marko  
Turku

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"A method to enable assessment of growth and death of micro-organisms"  
(Menetelmä mikro-organismien kasvun ja kuolemisen määrittämiseksi)

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*Eija Solja*

Eija Solja  
Apulaistarkastaja

Maksu 300,- mk  
Fee 300,- FIM

Osoite: Arkadiankatu 6 A Puhelin: 09 6939 500 Telefax: 09 6939 5328  
P.O.Box 1160 Telephone: + 358 9 6939 500 Telefax: + 358 9 6939 5328  
FIN-00101 Helsinki, FINLAND

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## A METHOD TO ENABLE ASSESSMENT OF GROWTH AND DEATH OF MICRO-ORGANISMS

This invention relates to a method to enable the assessment of growth and death of a micro-organism within a chosen time period in an environment of interest.

### 5 BACKGROUND OF THE INVENTION

When studying growth and death of a micro-organism under the influence of specific environments, e.g. production and storage environments that e.g. could or could not be refrigerated, or involving chemicals or matrixes, e.g. antibiotics, microbial toxins, heavy metals and serum complement, microbial cultures are most often incubated for hours or days. In these circumstances death and growth occur simultaneously. If additionally some of the cells lyse, e.g. when analysing the serum complement, it is difficult to know to what one should compare the amount of living cells at the end of the experiment. Convenient methods to determine the number of living cells, e.g. by measuring luciferase bioluminescence, are known but if no more information is available it is impossible to assess to what extent growth or/and death of the micro-organisms takes or has taken place.

Growth rates and death rates of micro-organisms in specific environments are of interest in many areas. Death rates and growth rates of micro-organisms and especially harmful and/or pathogenic micro-organisms are of importance in risk assessments of products of the pharmaceutical industry and products for human consumption with regard to there production, storage and distribution to the consumers. Knowledge of death and growth rates of micro-organisms are of particular importance in specific applications such as in the development of

antibiotics, disinfectants and bactericidal products or monitoring of sterilisation, disinfection and cleaning processes.

Reporter genes coding for luminescent or/and fluorescent products have been introduced to micro-organisms to enable the assessment of the quantity or survival of living micro-organisms. Even simultaneous use of luminescent and fluorescent markers have been used (Fratamico et al., Journal of Food Protection, Vol 50 No 10, 1997, 1167-1173). Luminescent and fluorescent markers have, however, only been used as markers for survival of micro-organisms and the use of two different markers within one micro-organism enabling the differentiation between growth and death rates has not been reported.

#### OBJECT AND SUMMARY OF THE INVENTION

The object of the present invention is to provide a method to enable the assessment of the growth and death of a micro-organism within a chosen time period in an environment of interest. The method is characterised in that

- 15    a) at least two reporter genes are introduced to said micro-organism, wherein the reporter genes used code for luminescent and/or fluorescent products and within said time period and environment at least two said products of the following are produced:
  - i) an essentially stable product produced in a, within the environment of interest, essentially known proportion to the total amount of cells of said micro-organism that are or have been alive within said chosen time period,
  - 20    ii) a product present in said environment of interest in an essentially known proportion to the amount of cells alive at any moment within said chosen time period and
- 25

iii) an essentially stable product produced in a within the environment of interest essentially known proportion to the total amount of cells of the said micro-organism that have died within said chosen time period and which products can be measured through their luminescence and/or fluorescence;

5 b) the said micro-organism is incubated within the environment of interest and said luminescence and/or fluorescence is detected after said chosen time periods and

c) the growth and death rate of the said micro-organism is assessed based on at least two of the following:

10 i) the known proportion of luminescence or fluorescence to the amount of cells alive after any said chosen time period,

15 ii) the known proportion of luminescence or fluorescence to the total amount of cells that are or have been alive within any said chosen time period and

15 iii) the known proportion of luminescence or fluorescence to the total amount of cells that have died within any said chosen time period.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows plasmid pGFP+<sup>luc\*</sup> including genes for both GFP and firefly luciferase.

20 Figure 2a-2f shows the sequence of plasmid pGFP+<sup>luc\*</sup>.

Figure 3 shows fluorescence during growth phase of *E. coli* with plasmid pGFP+<sup>luc\*</sup> at 30 °C as a function of the concentration of ethanol in the cell culture.

Figure 4 shows luminescence during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of ethanol in the cell culture.

Figure 5 shows the amount of living cells, i.e. colony forming units, according to plating during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of ethanol in the cell culture.

Figure 6 shows the percentage of live cells according to live/dead staining and flow cytometric analysis during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of ethanol in the cell culture.

Figure 7 shows fluorescence before and after incubation with serum complement during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of serum complement in the cell culture.

Figure 8 shows luminescence before and after incubation with serum complement during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of serum complement in the cell culture.

15 Figure 9 shows the percentage of living cells according to plating during growth phase of *E. coli* with plasmid pGFP+luc\* at 30 °C as a function of the concentration of serum complement in the cell culture.

#### DETAILED DESCRIPTION OF THE INVENTION

The method according to the present invention can be used to assess the growth and death rate of a micro-organism within a chosen time period in any particular environment of interest. The method is applicable if two different marker genes can be introduced to the micro-organism that code for luminescent and/or fluorescent products, and the products of these fulfil at least two of the following three criteria:

20 a) a said luminescent product luminesces or said fluorescent product fluoresces in

an essentially known proportion to the amount of cells of said micro-organism alive within said chosen time period ;

b) a said luminescent product luminesces or said fluorescent product fluoresces in an essentially known proportion to the amount of cells of said micro-organism that 5 are or have been alive within said chosen time period, and

c) a said luminescent product luminesces or said fluorescent product fluoresces in an essentially known proportion to the amount of cells of said micro-organism that have died within said chosen time period.

In the present application the concept "micro-organism" means any micro-organism 10 into which marker genes can be introduced so, that they will function according to the invention. "Micro-organism" can therefore stand for bacteria, yeast or fungi.

The concept of "introducing a marker gene into a micro-organism" means any method by which a marker gene can be made to function within the micro-organism according to the invention. One way of introducing marker genes into micro-15 organism is by constructing a recombinant strain. This can be done by transforming a strain with a plasmid including the marker genes. An alternative way to introduce reporter genes to bacteria is to utilise transposable elements. In this technique, reporter genes are inserted between insertion sequences in a delivery plasmid. The plasmid is then introduced to a cell by e.g. conjugation or transformation, and once 20 inside the cell, genes surrounded by the insertion sequences are integrated into bacterial chromosome. Integration is stable, i.e. there is no need for a selectable marker such as antibiotic resistance.

Assessment of the growth rate and death rate of a micro-organism can be of interest in many specific environments. Within pharmaceutical research the effect of 25 different drugs and candidates for drugs, e.g. antibiotics, on the survival of pathogenic, but also the beneficial micro-organisms of the gut, is of interest. Thus

the ultimate interest is in the behaviour of these micro-organisms in a physiological environment affected by drugs.

Another vast area where the possibility of assessing growth and death rate of specific micro-organisms is of interest is that of production, processing, storage and distribution of all products for human consumption. In this area the behaviour of pathogenic or potentially harmful micro-organisms in the different environments of the life cycle of these products is of special interest and involves many different aspects such as the influence of temperature, humidity or light and the possible use of preservatives etc.

5      10 Additionally growth and death rates of micro-organisms can be of interest for environmental evaluations e.g. when evaluating the effect of emissions into the environment.

Luminescent or fluorescent products coded by reporter genes in different embodiments of this invention can vary as long as their proportion to either the total 15 amount of cells alive, to cells that are or have been alive, or to cells that have died is essentially known. Growth and death rate can be assessed if two of the following: cells alive, cells that are or have been alive, or cells that have died can be determined. Thus luminescence and/or fluorescence measured can be e.g. of a product which is expressed e.g. constitutively or triggered by a specific phase 20 (e.g. replication or death) of the lifecycle of each cell, is stable or labile or which luminescence or fluorescence is dependant on a factor that relates e.g. to a specific phase of the lifecycle of each cell. Depending on the individual characteristics of said product—how produced, stable or labile, possible dependence of its luminescence or fluorescence of said factors etc.—the measured luminescence or 25 fluorescence can be in proportion to one of the three unknown of which two must be known to be able to assess the growth rate and death rate of said cells.

According to one specific embodiment of the invention assessment of the growth and death rate of an *Esherichia coli* strain under the influence of different chemicals or matrixes was enabled by constructing a recombinant strain, which expresses both luciferase and GFP. Alltogether the effect of a number of different chemicals and matrixes, such as CdCl<sub>2</sub>, ethanol, the antibiotics chloramphenicol, rifampicin, and tetracyclin, as well as serum complement on said recombinant *E. coli* strain was tested and found applicable.

The invention will be described in more detail by the following study in which the growth rate and death rate of a recombinant *Esherichia coli* strain, which expresses both luciferase and GFP, is assessed under the influence of ethanol or serum complement.

### **Summary of the study**

Genes for luciferase and green fluorescent protein have recently raised interest as reporter genes. Luciferase is an enzyme that produces luminescence in the presence of substrate luciferin, molecular oxygen and ATP. Green fluorescent protein (GFP), produces green fluorescence when excited with light. Many mutated forms of GFP have been introduced: some have different excitation and emission wavelenghts from the wild type and some mutants form more stable proteins at higher temperatures.

We constructed a recombinant strain of *E. coli*, which expresses both luciferase and GFP. In our construction we used a mutant of GFP which is more stable at temperatures over +30 °C and it matures quicker than the wild type. Luciferase was from North American firefly, *Photinus pyralis*.

The *E. coli* strain MC1061 was transformed with a plasmid including genes for both GFP and firefly luciferase. Figure 1 describes the plasmid in general and Figure 2 shows the sequence of the plasmid. Essential codings of the sequence are as follows:

lac promoter	95-199
GFP	289-1008
firefly luciferase	1044-2696
$\beta$ -lactamase	3251-4111

In our construct, see Figure 1, the luciferase gene is situated next to the GFP gene and both genes are transcribed in the same direction. The transcription is started at the lac promoter in front of GFP. The lac promoter is constitutively active, because the MC1061 cells lack its repressor. The plasmid also has a gene for ampicillin resistance (beta-lactamase).

The transformed *E. coli* strain was propagated under the influence of different concentrations of ethanol or serum complement.

## Methods

### Growth conditions

One colony from a pure culture plate was inoculated in 5 ml of LB-medium with ampicillin (100  $\mu$ g/ml) and grown at +37 °C in a shaker, 250 rpm, for about three hours. After that, the number of cells per millilitre was determined with flow cytometry by using fluorescent spherical latex particles as a reference. One million cells were then removed to an erlenmeyer with 50 ml of LB medium and ampicillin. The culture was grown over night in a shaker, 190 rpm, at room temperature to prevent the culture from growing into the stationary phase during the night. In the morning, the culture was transferred to and grown in a shaker, 330 rpm, until the stationary phase was reached or used after growth at +30 °C for about 1 h to study the influence ethanol or serum complement as described below.

Influence of chemicals on the propagation of *E. coli*

The culture obtained as described above was used to study the influence of ethanol or serum complement as follows:

*Ethanol*

5 Ethanol (Aa, Primalco Oy) was diluted into pure water to obtain concentrations of 50, 45, 25, 10, 5, 1 and 0 % of ethanol when 500 µl of said dilution was added to 500 µl of said culture in an eppendorf tube. The mixture was vortexed and incubated for 5 minutes before measuring fluorescence and luminescence. Live cells were again counted by plating and also by live/dead staining. In the live/dead protocol 10 used the stain *cyto 9* stains all cells whereas propidium iodide stains only the dead cells. After staining, cells are passed through a flow cytometer, with which dead and live cells can be differentiated and their proportion determined. (Virta et al. (1998) Appl. Environ. Microbiol. 64: 515–519.)

*Serum complement*

15 The influence of serum complement on the said recombinant *E. coli* strain was studied using an incubation time of 90 min as described for a different recombinant *E. coli* strain used in Virta et al. (1998) Appl. Environ. Microbiol. 64: 515–519.

Fluorescence and luminescence measurements

20 The measurements were done with a combined fluoro- and luminometer, Fluoroscan Ascent FL, provided by Labsystems Ltd. (Helsinki, Finland). Cell growth was simultaneously followed with a flow cytometer.

For the measurements, 100 µl of bacterial culture was pipetted into the microtiter plate wells. Fluorescence was measured using 485 nm for excitation and 510 nm for emission. Measuring time was 20 ms. After the fluorescence measurement 100 µl of

luciferin in 0.1 M citric acid-sodium citrate buffer (pH 5.0) was dispensed into the wells and the plate was shaken for two minutes (shaking diameter 1 mm, 1 020 rpm), after which luminescence was recorded with a measuring time of 1000 ms.

### Plating

Samples for plating were diluted 10<sup>2</sup> to 10<sup>7</sup> fold with 150 mM NaCl and plated onto L agar plates (L broth containing 1.6 % agar). Colonies were counted after overnight incubation at 37 °C.

### Live/dead staining and Flow cytometric analysis

Bacteria from 1 000 µl of cell culture were used for live/dead staining and flow cytometric analysis using a LIVE/DEAD BacLight bacterial viability kit (catalogue no. L-7005) for microscopy and quantitative analysis obtained from Molecular Probes Europe (Leiden, The Netherlands) and Fluoresbrite beads (diameter, 1.8 µm) obtained from Polysciences Inc. (Warrington, Pa.) as described in Virta et al. (1998) *Appl. Environ. Microbiol.* 64: 515–519.

### **Results**

When the cultures were transferred to +30 °C, the cells grew logarithmically for 1—4 hours depending on the initial cell concentration. Luminescence and fluorescence rose logarithmically and were essentially constant per cell. Thus cell number could be assessed based on luminescence or fluorescence.

When ethanol was added in different concentrations to the growth medium (see Figures 5 and 6) death was, after a very short incubation period of 5 min, more or less unsignificant at ethanol concentrations below 5 % and became more significant

with increasing ethanol concentration reaching very pronounced significance at ethanol concentrations above 10 %. Correspondingly fluorescence (Figure 3) was essentially constant whatever the ethanol concentration in spite of dramatically decreasing corresponding live cell count according to plate count (Figure 5) and percentage of live cells according to the live/dead staining (Figure 6) whereas luminescence dropped dramatically essentially corresponding to the dramatic drop in plate count (Figure 5) and the percentage of live cells (Figure 6) with increased ethanol concentration.

The effect of serum complement on the growth and death of *E. coli* is shown in Figures 7 to 9. Fluorescence (Figure 7) and luminescence (Figure 8) are shown before (squares) and after (circles) incubation for 90 minutes with serum complement. Fluorescence (Figure 7) is slightly increased, during incubation regardless of the concentration of serum, whereas luminescence (Figure 8) decreases during incubation with increasing serum concentration. The decrease of luminescence during incubation with increasing concentrations of serum correlates clearly with the percentage of cells alive after incubation (Figure 9).

## CLAIMS

1. A method to enable the assessment of the growth rate and death rate of a micro-organism within a chosen time period in an environment of interest characterised in that
- 5    a) at least two reporter genes are introduced into said micro-organism, wherein the reporter genes used code for luminescent and/or fluorescent products and within said time period and environment at least two said products of the following are produced:
  - i) an essentially stable product produced in a, within the environment of interest, essentially known proportion to the total amount of cells of said micro-organism that are or have been alive within said chosen time period,
  - 10    ii) a product present in said environment of interest in an essentially known proportion to the amount of cells alive at any moment within said chosen time period, and
  - 15    iii) an essentially stable product produced in a, within the environment of interest, essentially known proportion to the total amount of cells of the said micro-organism that have died within said chosen time period, and which products can be measured through their luminescence and/or fluorescence;
- 20    b) the said micro-organism is incubated within the environment of interest and said luminescence and/or fluorescence is detected after said chosen time period, and
- 25    c) the growth and death rate of the said micro-organism is assessed based on at least two of the following:

- i) the known proportion of luminescence or fluorescence to the amount of cells alive after any said chosen time period,
- ii) the known proportion of luminescence or fluorescence to the total amount of cells that are or have been alive within any said chosen time period, and
- iii) the known proportion of luminescence or fluorescence to the total amount of cells that have died within any said chosen time period.

5

2. The method according to claim 1 **characterised** in that said micro-organism is a gram negative bacteria, e.g. *Escherichia coli*.

10 3. The method according to claim 1 or 2 **characterised** in that

- a) one reporter gene coding for a luminescent product is luciferase, which is used for the determination of amount of cells alive at any moment within said chosen time period, and
- b) another reporter gene coding for a fluorescent product is green fluorescent

15 15 protein (GFP), which is used for the determination of total amount of cells of said micro-organism that are or have been alive within said chosen time period.

4. The method according to claim 1 or 2 **characterised** in that said reporter genes are introduced into said micro-organism in a plasmid.

5. A method according to the methods of claim 3 and 4 **characterised** in that said

20 20 plasmid is pGFP+*luc*\* (Figure 1 and 2a-2f).

## ABSTRACT

A method to enable the assessment of the growth rate and death rate of a micro-organism within a chosen time period in an environment of interest. The method is characterised in that

- a) two reporter genes are introduced to said micro-organism wherein, the reporter genes used code for luminescent and/or fluorescent products and at least two of the following products: an essentially stable product produced in an essentially known proportion to the total amount of cells of said micro-organism that are or have been alive within said chosen time period; a product present in an essentially known proportion to the amount of cells alive at any moment within said chosen time period; and an essentially stable product produced in an essentially known proportion to the total amount of cells of the said micro-organism that have died within said chosen time period, and said products can be measured through their luminescence and/or fluorescence;
- b) the said micro-organism is incubated and said luminescence and/or fluorescence is detected after said chosen time periods, and
- c) the growth and death rate of the said micro-organism is assessed based on at least two of the following: the known proportion of luminescence or fluorescence to the amount of cells alive after any said chosen time period; the known proportion of luminescence or fluorescence to the total amount of cells that are or have been alive within any said chosen time period; and the known proportion of luminescence or fluorescence to the total amount of cells that have died within any said chosen time period.

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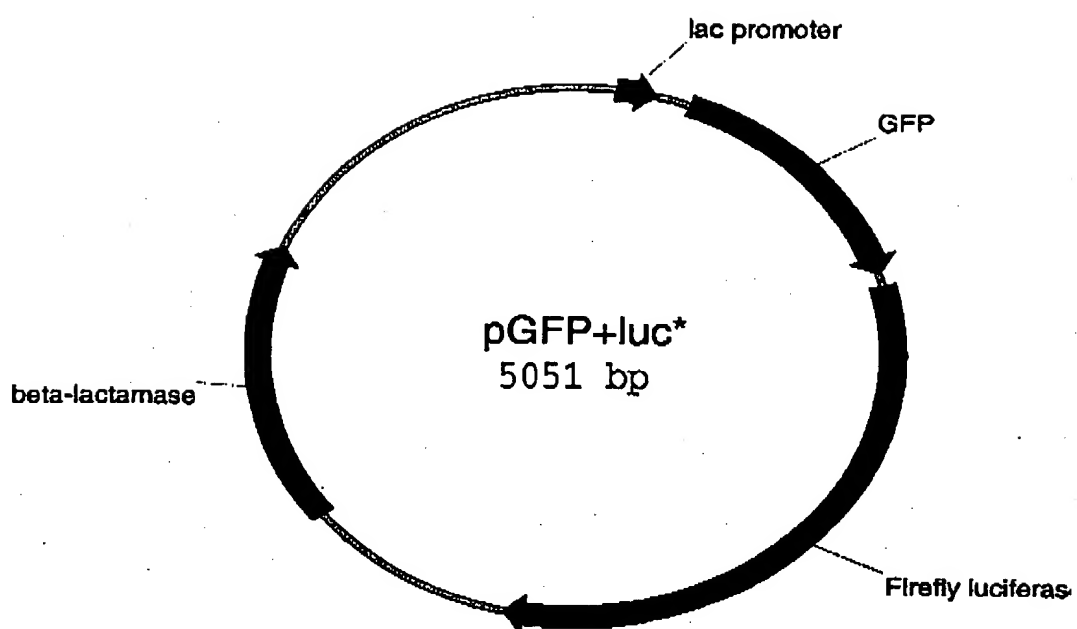


Figure 1

1 AGCGCCCAAT ACGCAAACCG CCTCTCCCCG CGCGTTGGCC GATTCAATTAA  
 TCGGGGTTA TGCCTTGGC GGAGAGGGGC GCGAACCGG CTAAGTAATT  
 51 TGCAGCTGGC ACGACAGGTT TCCCAGCTGG AAAGCGGGCA GTGAGCGCAA  
 ACGTCGACCG TGCTGTCCAA AGGGCTGACC TTTCGCCCCT CACTCGCGTT  
 101 CGCAATTAAT GTGAGTTAGC TCACTCATTA GGCACCCCCAG GCTTTACACT  
 GCGTTAATTA CACTCAATCG AGTGAGTAAT CGTGGGGTC CGAAATGTGA  
 151 TTATGCTTCC GGCTCGTATG TTGTGTGGAA TTGTGAGCGG ATAACAATT  
 AATAACGAAGG CCGAGCATAAC AACACACCTT AACACTCGCC TATTGTTAAA  
 201 CACACAGGAA ACAGCTATGA CCATGATTAC GCCAAGCTTG CATGCCTGCA  
 GTGTGTCCTT TGTCGATACT GGTACTAATG CGGTTCGAAC GTACGGACGT  
 251 GGTCGACTCT AGAGGATCCC CGGGTACCCGG TCGCCACCAC GGTGAGCAAG  
 CCAGCTGAGA TCTCCTAGGG GCCCATGGCC AGCGGTGGTA CCACTCGTTC  
 301 GGCGAGGAGC TGTCACCCGG GGTGGTGCCCG ATCCTGGTCC AGCTGGACGG  
 CCCGTCCTCG ACAAGTGGCC CCACCACGGG TAGGACCACG TCGACCTGCC  
 351 CGACGTAAAC GGCCACAAAGT TCAGCGTGTGCG CGGGCGAGGGC GAGGGCGATG  
 GCTGCATTTG CCGGTGTTCA AGTCGCACAG GCGCCTCCCG CTCCCGCTAC  
 401 CCACCTACGG CAAGCTGACC CTGAAGTTCA TCTGCACCAC CGGCAAGCTG  
 GGTGGATGCC GTTCGACTGG GACTTCAGT AGACGTGGTG GCGTTCGAC  
 451 CCCGTGCCCT GGCCCACCCCT CGTGACCACC CTGACCTACG GCGTGCAGTG  
 GGGCACGGGA CCGGGTGGGA GCACTGGTGG GACTGGATGC CGCACGTCAC  
 501 CTTCAAGCCGC TACCCCGACC ACATGAAGCA GCACGACTTC TTCAAGTCCG  
 GAAGTCGGCG ATGGGGCTGG TGTACTTCGT CGTGTGAAG AAGTCAGGC  
 551 CCATGCCCGA AGGCTACGTC CAGGAGCGCA CCATCTCTT CAAGGACGAC  
 GGTACGGGCT TCCGATGCA GTCCTCGCGT GGTAGAAGAA GTTCTGCTG  
 601 GGCAACTACA AGACCCCGCGC CGAGGTGAAG TTGAGGGCG ACACCCCTGGT  
 CCGTTGATGT TCTGGCGCG GCTCCACTTC AAGCTCCCGC TGTGGGACCA  
 651 GAACCGCATC GAGCTGAAGG GCATCGACTT CAAGGAGGAC GGCAACATCC  
 CTTGGCGTAG CTCGACTTCC CGTAGCTGAA GTTCTCCTG CGTTGTAGG  
 701 TGGGGCACAA GCTGGAGTAC AACTACAACA GCCACAACGT CTATATCATG  
 ACCCCGTGTT CGACCTCATG TTGATGTTGT CGGTGTTGCA GATATAGTAC  
 751 GCCGACAAGC AGAAGAACCGG CATCAAGGTG AACTTCAGA TCCGCCACAA  
 CGGCTGTTCG TCTTCTTGCC GTAGTTCCAC TTGAAGTTCT AGGCAGGTGTT  
 801 CATCGAGGAC GGCAGCGTGC AGCTGCCGA CCACCTACCG CAGAACACCC  
 GTAGCTCCTG CCGTCGCACG TCGAGCGGCT GGTGATGGTC GTCTTGTGGG  
 851 CCATCGGCAGA CGGCCCCGTG CTGCTGCCCG ACAACCACCA CCTGAGCACC  
 GGTAGCCGCT GCCGGGGCAC GACGACGGG TGTTGGTGT GGACTCGTGG  
 901 CAGTCCGCCCG TGACCAAAGA CCCCAACCGAG AAGCGCGATC ACATGGTCCT  
 GTCAGGGCGGG ACTCGTTCT GGGGTTGCTC TTCGCGCTAG TGTACCAAGGA  
 951 GCTGGAGTTC GTGACCGCCG CGGGGATCAC TCTCGGCATG GACGAGCTGT  
 CGACCTCAAG CACTGGCGGC GCCCCTAGTG AGAGCCGTAC CTGCTCGACA

Figure 2a

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1001 ACAAGTAAAG CGGCCGCTCT AGAACTAGTG GATCCCCGT ACCATGGAAG  
 TGTTCATTTG GCCGGCGAGA TCTTGATCAC CTAGGGGGCA TGGTACCTTC  
 1051 ACGCCAAAAA CATAAAAGAAA GGCCCGGGCGC CATTCTATCC GCTAGAGGAT  
 TGC GGTTTT GTATTTCTTT CCGGGCCCGC GTAAGATAGG CGATCTCCTA  
 1101 GGAACCGCTG GAGAGCAACT GCATAAGGCT ATGAAGAGAT ACGCCCTGGT  
 CCTTGGCGAC CTCTCGTTGA CGTATTCCGA TACTTCTCTA TGC GGGACCA  
 1151 TCCTGGAACA ATTGCTTTA CAGATGCACA TATCGAGGTG AACATCACGT  
 AGGACCTTGT TAACGAAAAT GTCTACGTGT ATAGCTCCAC TTGTAGTGCA  
 1201 ACGCGGAATA CTTCGAAATG TCCGTTCGGT TGGCAGAAC TATGAAACGA  
 TGC GCGCTTAT GAAGCTTTAC AGGCAAGCCA ACCGTCTTCG ATACTTTGCT  
 1251 TATGGGCTGA ATACAAATCA CAGAATCGTC GTATGCAGTG AAAACTCTCT  
 ATACCCGACT TATGTTAGT GTCTTAGCAG CATACTCAC TTTTGAGAGA  
 1301 TCAATTCTTT ATGCCGGTGT TGGCGCGTT ATTTATCGGA GTTGCAGTTG  
 AGTTAAGAAA TACGGCCACA ACCCGCGCAA TAAATAGCCT CAACGTCAAC  
 1351 CGCCCGCGAA CGACATTTAT AATGAACGTG AATTGCTCAA CAGTATGAAC  
 CGGGCGCTT GCTGTAAATA TTACTGCAC TAAACGAGTT GTCATACTTG  
 1401 ATTTCGCAGC CTACCGTAGT GTTGTTCAC AAAAAGGGGT TGCAAAAAT  
 TAAAGCGTCG GATGGCATCA CAAACAAAGG TTTTCCCCA ACAGTTTTTA  
 1451 TTTGAACGTG CAAAAAAAT TACCAATAAT CCAGAAAATT ATTATCATGG  
 AAAC TTGCAC GTTTTTTTA ATGTTATTA GGTCTTTAA TAATAGTACC  
 1501 ATTCTAAAAC GGATTACCAAG GGATTTCACT CGATGTACAC GTTCGTACAC  
 TAAGATTTG CCTAATGGTC CCTAAAGTCA GCTACATGTG CAAGCAGTGT  
 1551 TCTCATCTAC CTCCCGGTTT TAATGAATAC GATTTGTAC CAGAGTCCTT  
 AGAGTAGATG GAGGGCCAAA ATTACTTATG CTAAAACATG GTCTCAGGAA  
 1601 TGATCGTGAC AAAACAATTG CACTGATAAT GAACTCCTCT GGATCTACTG  
 ACTAGCACTG TTTGTTAAC GTGACTATTA CTTGAGGAGA CCTAGATGAC  
 1651 GGTTACCTAA GGGTGTGGCC CTTCCGCATA GAACTGCCTG CGTCAGATTC  
 CCAATGGATT CCCACACCGG GAAGGGCGTAT CTTGACGGAC GCAGTCTAAG  
 1701 TCGCATGCCA GAGATCCTAT TTTGGCAAT CAAATCATTC CGGATACTGC  
 AGCGTACGGT CTCTAGGATA AAAACC GTTAAAG GCCTATGACG  
 1751 GATTTTAAGT GTTGTCCAT TCCATCACGG TTTGGAATG TTTACTACAC  
 CTAAAATTCA CAACAAGGTA AGGTAGTGCG AAAACCTTAC AAATGATGTG  
 1801 TCGGATATTT GATATGTGGA TTTCGAGTCG TCTTAATGTA TAGATTGAA  
 AGCCTATAAA CTATACACCT AAAGCTCAGC AGAATTACAT ATCTAAACTT  
 1851 GAAGAGCTGT TTTTACGATC CCTTCAGGAT TACAAAATTG AAAGTGCCTT  
 CTTCTCGACA AAAATGCTAG GGAAGTCCTA ATGTTTTAAG TTTCACGCAA  
 1901 GCTAGTACCA ACCCTATTTT CATTCTTCGC CAAAAGCACT CTGATTGACA  
 CGATCATGGT TGGGATAAAA GTAAGAAGCG GTTTCGTGA GACTAACTGT  
 1951 AATACGATTT ATCTAATTCA CACGAAAATTG CTTCTGGGGG CGCACCTCTT  
 TTATGCTAAA TAGATTAAT GTGCTTTAAC GAAGACCCCC CGGTGGAGAA

Figure 2b

2001 TCGAAAGAAG TCGGGGAAGC GGTTGCAAAA CGCTTCCATC TTCCAGGGAT  
 AGCTTCTTC AGCCCCCTCG CCAACGTTT GCGAAGGTAG AAGGTCCCTA  
 2051 ACGACAAGGA TATGGGCTCA CTGAGACTAC ATCAGCTATT CTGATTACAC  
 TGCTGTTCT ATACCCGAGT GACTCTGATG TAGTCGATAA GACTAATGTG  
 2101 CCGAGGGGGGA TGATAAACCG GGCAGCGGTGCG GTAAAGTTGT TCCATTCTT  
 GGCTCCCCCT ACTATTGGC CCGCGCCAGC CATTCAACA AGGTAAAAAA  
 2151 GAAGCGAAGG TTGTGGATCT GGATACCGGG AAAACGCTGG GCGTTAACCA  
 CTTCGCTTCC AACACCTAGA CCTATGGCCC TTTTGCACCG CGCAATTAGT  
 2201 GAGAGGCAGA TTATGTGTCA GAGGACCTAT GATTATGTCC GGTTATGTAA  
 CTCTCCGCTT AATACACAGT CTCCCTGGATA CTAATACAGG CCAATACATT  
 2251 ACAATCCGGA AGCGACCAAC GCCTTGATG ACAAGGATGG ATGGCTACAT  
 TGTAGGCCT TCGCTGGTT CGGAACTAAC TGTTCCCTACC TACCGATGTA  
 2301 TCTGGAGACA TAGCTTACTG GGACGAAGAC GAACACTTCT TCATAGTTGA  
 AGACCTCTGT ATCGAAATGAC CCTGCTTCTG CTTGTGAAGA AGTATCAACT  
 2351 CCGCTTGAAG TCTTTAATTAA AATACAAAGG ATACCAGGTG GCCCCCGCTG  
 GGCAGACTTC AGAAATTAAAT TTATGTTCC TATGGTCCAC CGGGGGCGAC  
 2401 AATTGGAGTC GATATTGTTA CAACACCCCA ACATCTTCGA CGCGGGCGTG  
 TTAACCTCAG CTATAACAAT GTTGTGGGGT TGTAGAAGCT CGGCCCGCAC  
 2451 GCAGGTCTTC CCGACGATGA CGCCGGTGAA CTTCCCGCCG CCGTTGTTGT  
 CGTCCAGAAG GGCTGCTACT GCGGCCACTT GAAGGGCGGC GGCAACAACA  
 2501 TTTGGAGCAC GGAAAGACGA TGACGGAAAA AGAGATCGTG GATTACGTG  
 AAACCTCGTG CCTTTCTGCT ACTGCCTTT TCTCTAGCAC CTAATGCAGC  
 2551 CCAGTCAGT AACAAACCGCC AAAAAGTTGC GCGGAGGAGT TGTGTTGTG  
 GGTCAAGTCA TTGTTGGCGG TTTTCAACG CGCCTCCTCA ACACAAACAC  
 2601 GACGAAGTAC CGAAAGGTCT TACCGGAAAA CTCGACGCCA GAAAATCAG  
 CTGCTTCATG GCTTCCAGA ATGGCCTTT GAGCTGCCTT CTTTTAGTC  
 2651 AGAGATCCTC ATAAAGGCCA AGAAGGGCGG AAAGTCCAAA TTGTAAAATG  
 TCTCTAGGAG TATTCCGGT CCTTCCCGCC TTTCAGGTTT AACATTTTAC  
 2701 TAACTGTATT CAGCGATGAC GAAATTCTTA GCTATTGTAA TACTCTAGGG  
 ATTGACATAA GTCGCTACTG CTTTAAGAAT CGATAACATT ATGAGATCCC  
 2751 GCTGCAGGAA TTGATATCA AGCTTATCGA TACCGTCGAC CTCGAGGGGG  
 CGACGTCCTT AAGCTATAGT TCGAATAGCT ATGGCAGCTG GAGCTCCCC  
 2801 GGCCCTTCG TCTCGCGCGT TTGCGGTGATG ACGGTGAAAA CCTCTGACAC  
 CCGGGAAAGC AGAGCGCGA AAGCCACTAC TGCCACTTT GGAGACTGTG  
 2851 ATGCAGCTCC CGGAGACGGT CACAGCTTGT CTGTAAGCGG ATGCCGGGAG  
 TACGTCGAGG GCCTCTGCCA GTGTCGAACA GACATTGCC TACGGCCCTC  
 2901 CAGACAAGCC CGTCAGGGCG CGTCAGCGGG TGTTGGCGGG TGTCGGGGCT  
 GTCTGTTCGG GCAGTCCCGC GCAGTCGCC ACAACCGCCC ACAGCCCCGA  
 2951 GGCTTAACCA TGCGGCATCA GAGCAGATTG TACTGAGAGT GCACCATATG  
 CCGAATTGAT ACGCCGTAGT CTCGTCTAAC ATGACTCTCA CGTGGTATAC

Figure 2c

3001 CGGTGTGAAA TACCGCACAG ATGCGTAAGG AGAAAATACC GCATCAGGCG  
 GCCACACTT ATGGCGTGTC TACGCATTCC TCTTTATGG CGTAGTCCGC  
 3051 GCCTTAAGGG CCTCGTGATA CGCCTATTT TATAGGTTAA TGTCAATGATA  
 CGGAATTCCC GGAGCACTAT GCGGATAAAA ATATCCAATT ACAGTACTAT  
 3101 ATAATGGTTT CTTAGACGTC AGGTGGCACT TTTGGGGAA ATGTGCGCGG  
 TATTACCAAA GAATCTGCAG TCCACCGTGA AAAGCCCCTT TACACGCGCC  
 3151 AACCCCTATT TGTATTTT TCTAAATACA TTCAAATATG TATCCGCTCA  
 TTGGGGATAA ACAAAATAAAA AGATTTATGT AAGTTTATAC ATAGGGCGAGT  
 3201 TGAGACAATA ACCCTGATAA ATGCTTCAT AATATTGAAA AAGGAAGAGT  
 ACTCTGTTAT TGGGACTATT TACGAAGTTA TTATAACTTT TTCCCTCTCA  
 3251 ATGAGTATTG AACATTCCG TGTCGCCCTT ATTCCCTTTT TTGCGGCATT  
 TACTCATAAG TTGTAAAGGC ACAGCGGGAA TAAGGGAAA AACGCCGTAA  
 3301 TTGCCTTCCT GTTTTGCTC ACCCAGAAAC GCTGGTGAAA GTAAAAGATG  
 AACGGAAGGA CAAAAACGAG TGGGTCTTTG CGACCACCTT CATTTCCTAC  
 3351 CTGAAGATCA GTTGGGTGCA CGAGTGGGTT ACATCGAACT GGATCTCAAC  
 GACTCTAGT CAACCCACGT GCTCACCCAA TGTAGCTTGA CCTAGAGTTG  
 3401 AGCGGTAAGA TCCTTGAGAG TTTTCGCCCG GAAGAACGTT TTCCAATGAT  
 TCGCCATTCT AGGAACCTCTC AAAAGCGGGG CTTCTTGCAA AAGGTTACTA  
 3451 GAGCACTTTT AAAGTTCTGC TATGTGGCGC GGTATTATCC CGTATTGACG  
 CTCGTAAAAA TTTCAAGACG ATACACCGCG CCATAATAGG GCATAACTGTC  
 3501 CCGGGCAAGA GCAACTCGGT CGCCGCATAC ACTATTCTCA GAATGACTTG  
 GGCCCCTCT CGTTGAGCCA GCGCGTATG TGATAAGAGT CTTACTGAAC  
 3551 GTTGAGTACT CACCAGTCAC AGAAAAGCAT CTTACGGATG GCATGACAGT  
 CAACTCATGA GTGGTCAGTG TCTTTTCGTA GAATGCCCTAC CGTACTGTCA  
 3601 AAGAGAATTG TGCACTGCTG CCATAACCAT GAGTGATAAC ACTGCGGCCA  
 TTCTCTTAAT ACGTCAACGAC GGTATTGGTA CTCACTATTG TGACGCCGGT  
 3651 ACTTACTTCT GACAACGATC GGAGGACCGA AGGAGCTAAC CGCTTTTTG  
 TGAATGAAGA CTGTTGCTAG CCTCCTGGCT TCCTCGATTG GCGAAAAAAC  
 3701 CACAACATGG GGGATCATGT AACTCGCCTT GATCGTTGGG AACCGGAGCT  
 GTGGTGTACC CCCTAGTACA TTGAGCGGAA CTAGCAACCC TTGGCCTCGA  
 3751 GAATGAAGCC ATACCAAACG ACGAGCGTGA CACCACGATG CCTGTAGCAA  
 CTTACTTCGG TATGGTTGTC TGCTCGCACT GTGGTGCTAC GGACATCGTT  
 3801 TGGCAACAAAC GTTGGCCTAA CTATTAACG GCGAACTACT TACTCTAGCT  
 ACCGTTGTTG CAACCGTGT GATAATTGAC CGCTTGATGA ATGAGATCGA  
 3851 TCCCGGCAAC AATTAATAGA CTGGATGGAG GCGGATAAAAG TTGCAGGACC  
 AGGGCCGTTG TTAATTATCT GACCTACCTC CGCCTATTTC AACGTCCTGG  
 3901 ACTTCTGCGC TCGGCCCTTC CGGCTGGCTG GTTTATTGCT GATAAAATCTG  
 TGAAGACGCC AGCCGGGAAG GCCGACCGAC CAAATAACGA CTATTTAGAC  
 3951 GAGCCGGTGA GCGTGGGTCT CGCGGTATCA TTGCAGCACT GGGGCCAGAT  
 CTCGGCCACT CGCACCCAGA GCGCCATAGT AACGTCGTGA CCCCCGGTCTA

Figure 2d

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4001 GGTAAAGCCCT CCCGTATCGT AGTTATCTAC ACGACGGGGA GTCAGGCAAC  
 CCATTGGGA GGGCATAGCA TCAATAGATG TGCTGCCCT CAGTCGTTG  
 4051 TATGGATGAA CGAAATAGAC AGATCGCTGA GATAGGTGCC TCACTGATTA  
 ATACCTACTT GCTTTATCTG TCTAGCGACT CTATCCACGG AGTGAAC  
 4101 AGCATTGGTA ACTGTCAAGAC CAAGTTACT CATATATACT TTAGATTGAT  
 TCGTAACCAT TGACAGTCTG GTTCAAATGA GTATATATGA AATCTAACTA  
 4151 TAAAAACTTC ATTTTTAATT TAAAAGGATC TAGGTGAAGA TCCTTTGGA  
 AATTTGAAG TAAAAATTAA ATTTTCCTAG ATCCACTTCT AGGAAAAACT  
 4201 TAATCTCATG ACCAAAATCC CTTAACGTGA GTTTTCGTTC CACTGAGCGT  
 ATTAGAGTAC TGGTTTTAGG GAATTGCACT CAAAAGCAAG GTGACTCGCA  
 4251 CAGACCCCGT AGAAAAGATC AAAGGATCTT CTTGAGATCC TTTTTTCTG  
 GTCTGGGCA TCTTTCTAG TTTCCTAGAA GAACTCTAGG AAAAAAAGAC  
 4301 CGCGTAATCT GCTGCTTGCA AACAAAAAAA CCACCGCTAC CAGCGGTGGT  
 GCCCATAGA CGACGAACGT TTGTTTTTT GGTGGCGATG GTCGCCACCA  
 4351 TTGTTGCCG GATCAAGAGC TACCAACTCT TTTTCCGAAG GAACTGGCT  
 AACAAACGGC CTAGTTCTCG ATGGTTGAGA AAAAGGCTTC CATTGACCGA  
 4401 TCAGCAGAGC GCAGATACCA AATACTGTCC TTCTAGTGTAA GCGTAGTTA  
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 4451 GGCCACCACT TCAAGAACTC TGTAGCACCG CCTACATACC TCGCTCTGCT  
 CCGGTGGTGA AGTTCTTGAG ACATCGTGGC GGATGTATGG AGCGAGACGA  
 4501 AATCCTGTTA CCAGTGGCTG CTGCCAGTGG CGATAAGTCG TGTCTTACCG  
 TTAGGACAAT GGTCACCGAC GACGGTCACC GCTATTCAAG ACAGAATGGC  
 4551 GGTTGGACTC AAGACGATAG TTACCGGATA AGGCCGAGCG GTCGGGCTGA  
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 4601 ACGGGGGGTT CGTGCACACA GCCCAGCTTG GAGCGAACGA CCTACACCGA  
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 4651 ACTGAGATAC CTACAGCGTG AGCTATGAGA AAGCGCCACG CTTCCCGAAG  
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 4701 GGAGAAAGGC GGACAGGTAT CCGGTAAGCG GCAGGGTCGG AACAGGAGAG  
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 4751 CGCACGAGGG AGCTTCCAGG GGGAAACGCC TGTTATCTT ATAGTCCTGT  
 GCGTGTCTCC TCGAAGGTCC CCCCTTGCAG ACCATAGAAA TATCAGGACA  
 4801 CGGGTTTCGC CACCTCTGAC TTGAGCGTCG ATTTTGTGA TGCTCGTCAG  
 GCCCAAAGCG GTGGAGACTG AACTCGCAGC TAAAAACACT ACGAGCAGTC  
 4851 GGGGGCGGAG CCTATGGAAA AACGCCAGCA ACGCGGCCTT TTTACGGTTC  
 CCCCCGCCTC GGATACCTT TTGCGGGTGT TGCGCCGGAA AAATGCCAAG  
 4901 CTGGCCTTT GCTGGCCTT TGCTCACATG TTCTTCCCTG CGTTATCCCC  
 GACCGGAAAA CGACCGGAAA ACGAGTGTAC AAGAAAGGAC GCAATAGGGG  
 4951 TGATTCTGTG GATAACCGTA TTACCGCCTT TGAGTGTGAGCT GATACCGCTC  
 ACTAAGACAC CTATTGCCAT AATGGCGGAA ACTCACTCGA CTATGGCGAG

Figure 2e

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5001 GCCGCAGCCG AACGACCGAG CGCAGCGAGT CAGTGAGCGA GGAAGCGGAA  
CGGCCTCGGC TTGCTGGCTC GCGTCGCTCA GTCACTCGCT CCTTCGCCCTT

5051 G  
C

Figure 2f

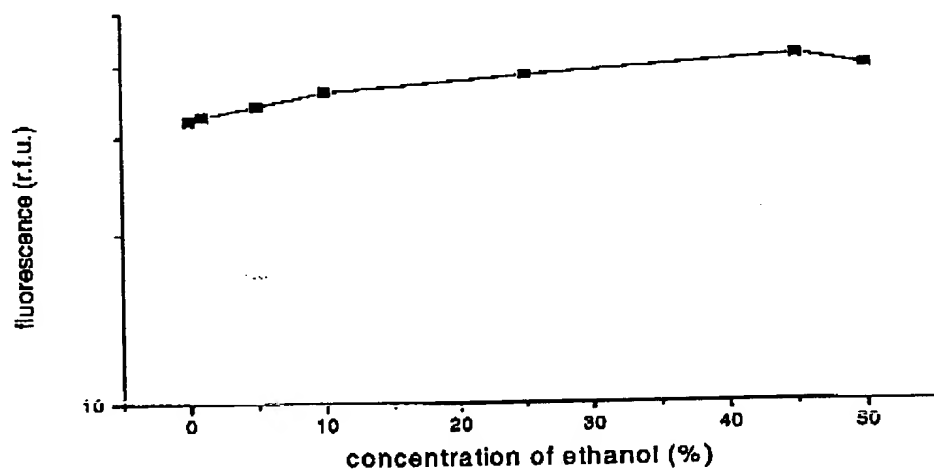


Figure 3

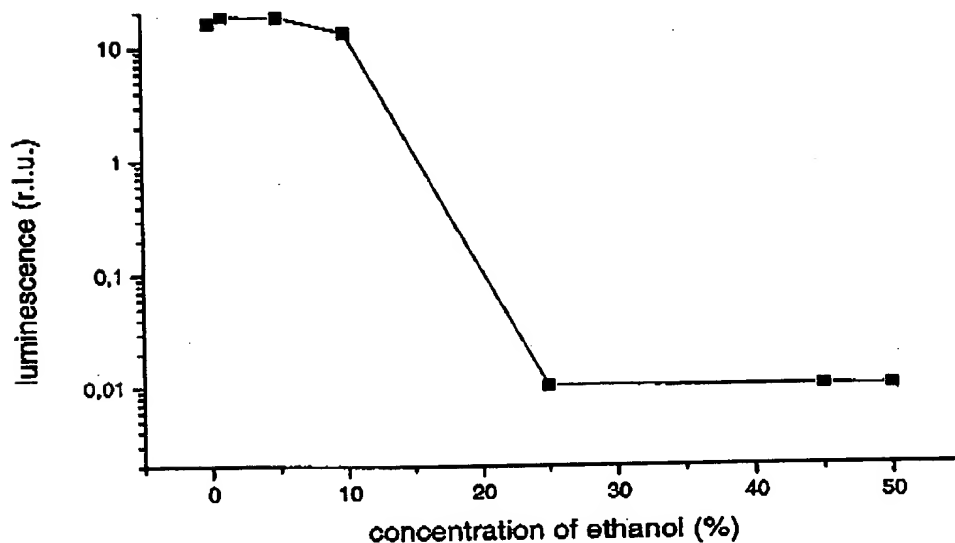


Figure 4

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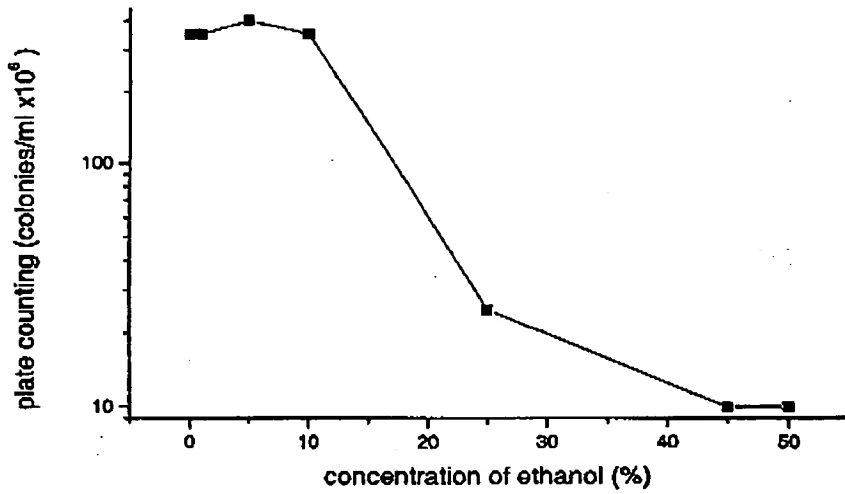


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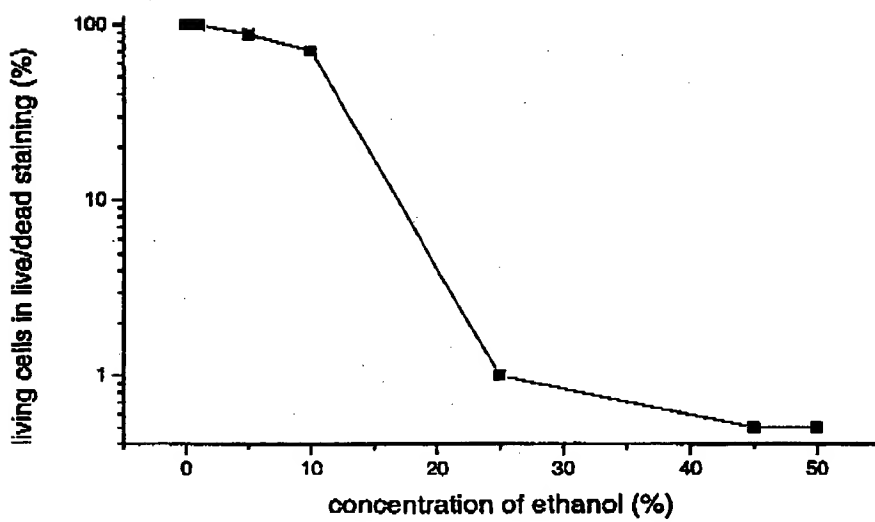


Figure 6

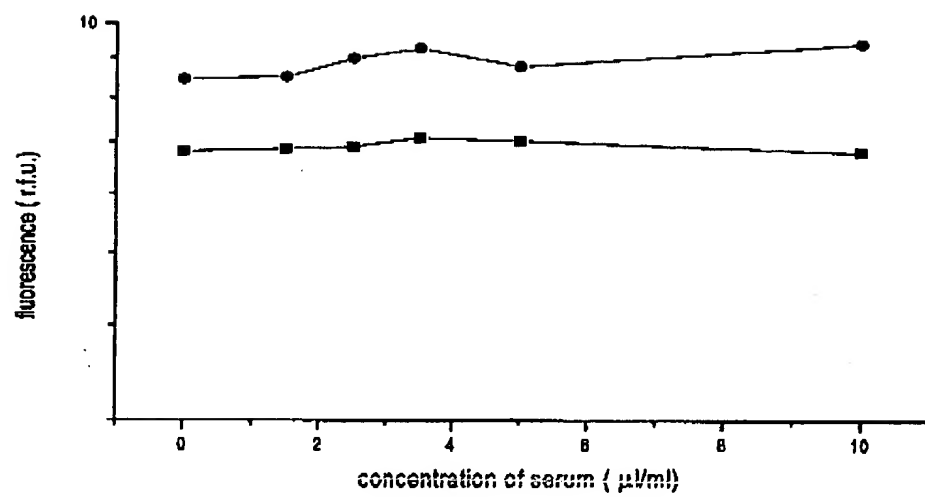


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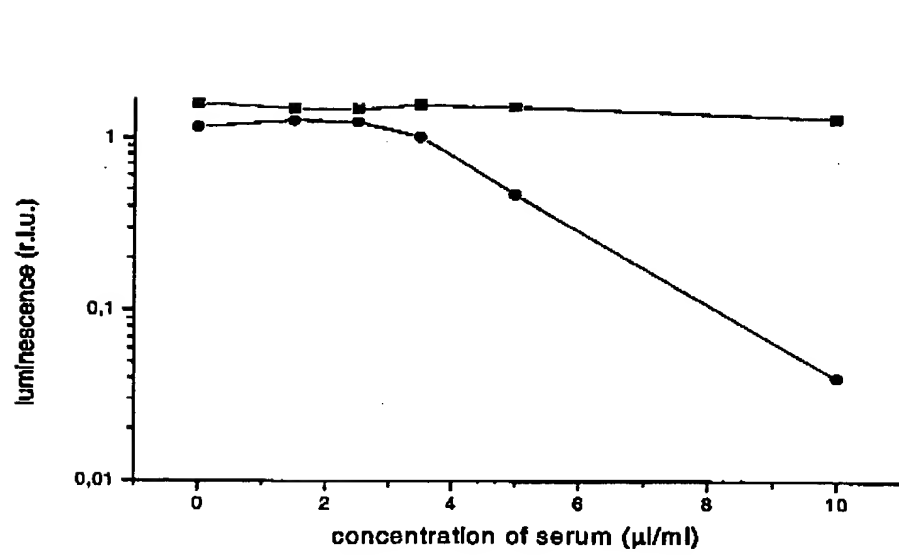


Figure 8

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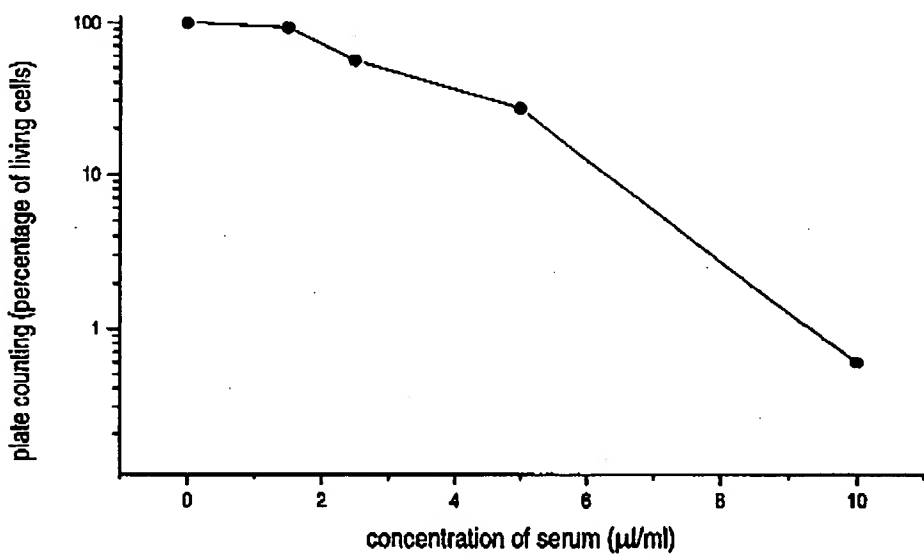


Figure 9